(FILE 'HOME'	ENTERED	AT	16:36:46	on	25	JAN	2007)	
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60/676882 Search by Eilhice 1/25/2007

	FILE	'REGISTRY'	ENTERED	AT	16:37:40	ON	25	JAN	2007
L1		1 S 3-1	HYDROXYPI	ROPA	NAL/CN				

1 S 1,3-PROPANEDIOL/CN

L3 1 S SODIUM HYDROXIDE/CN

L2

FILE 'CAPLUS, CAOLD' ENTERED AT 16:39:45 ON 25 JAN 2007

	LIDE	CAPIO	J	, C	מעטסר	ENTERED
L4		137	S	L1	AND	L2
L5		4	S	L4	AND	L3
L6		1	S	L5	AND	VISCO?
L7		11	S	L4	AND	HYDROXIDE
L8		7	s	L7	NOT	L5
L9		7	S	L8	AND	HYDROGEN?
L10		0	S	L9	AND	VISCO?

ANSWER 1 OF 4 CAPLUS COPYRIGHT 2007 ACS ON STN

2006:491425 CAPLUS

144:489080

Catalyst for the production of 1,3-propanediol by catalytic hydrogenation

of 3-bydroxymropanal L5 ANDN ΤI of 3-hydroxypropanal IN Eng, John Harvey PA USA U.S. Pat. Appl. Publ., 13 pp. SO CODEN: USXXCO DTPatent English LA FAN.CNT 1 DATE PATENT NO. KIND APPLICATION NO. DATE _____ _ - - -_____ _____ _____ US 2006111595 20060525 US 2004-996756 20041124 PRAI US 2004-996756 20041124 A process for preparing a catalyst, comprising sequentially: (a) saturating a TiO2 support with aqueous ruthenium solution; (b) treating the ruthenium saturated TiO2 support with an aqueous base solution; and (c) curing the ruthenium saturated TiO2 to impregnate the TiO2 with ruthenium. Preferably, subsequent to step (c) the process comprises sequentially (d) washing the ruthenium impregnated TiO2 with water, (e) drying the washed ruthenium impregnated TiO2, and (f) heat treating the ruthenium impregnated TiO2 under reducing conditions at temps. of 150-800° C. A catalyst prepared by the process. A process for the production of 1,3-propanediol, comprising: (a) providing 3-hydroxypropanal and the catalyst, and (b) hydrogenating the 3-hydroxypropanal in the presence of the catalyst. ANSWER 2 OF 4 CAPLUS COPYRIGHT 2007 ACS on STN L5 2004:308390 CAPLUS AN140:323182 DN Reduction of the viscosity of reactive heavy byproducts during the ΤI production of 1,3-propanediol using other than sodium hydroxide James, Talmadge Gail; Komplin, Glenn Charles; Nielsen, Edward Lewis; IN Knifton, John Frederick; Powell, Joseph Broun; Weider, Paul Richard Shell Oil Company, USA PΑ so PCT Int. Appl., 15 pp. CODEN: PIXXD2 DTPatent LA English FAN.CNT 1 KIND APPLICATION NO. PATENT NO. DATE DATE --------------------_____ WO 2003-US31216 PΙ WO 2004031108 A1 20040415 20031002 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG

US 2003-676682

CA 2003-2500615

AU 2003-279749

EP 2003-773090

AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK

20031001

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US 2005043570

AU 2003279749

CA 2500615

EP 1546075

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20040415

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20050629

	BR	2003015025	A	20050816	BR	2003-15025	20031002
	CN	1703388	A	20051130	CN	2003-80100866	20031002
	JP	2006502202	T	20060119	JP	2004-542042	20031002
PRAI	US	2002-415676P	P	20021003			
	WO	2003-US31216	W	20031002			

AB An improvement upon the process for the production of 1,3-propanediol is presented where an aqueous solution of 3-hydroxypropanal is formed, the catalyst,

if any, used in the formation is removed from the solution, sodium hydroxide is added to the solution to neutralize any acid therein such that the pH is ≥5, the neutralized aqueous solution is subjected to hydrogenation to produce a crude 1,3-propanediol mixture which is distilled to produce 1,3-propanediol, water, and reactive heavy components. The improvement comprises replacing the sodium hydroxide with a hydroxide selected from the group consisting of ammonium hydroxide, alkali metal hydroxides other than sodium hydroxide, and alkaline earth metal hydroxides to reduce the viscosity of the reactive heavy components.

RE.CNT 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L5 ANSWER 3 OF 4 CAPLUS COPYRIGHT 2007 ACS on STN
- AN 1999:177140 CAPLUS
- DN 130:330016
- TI In-situ infrared reflectance spectroscopic study of propanediol electrooxidation at platinum and gold Part 1: 1,3-propanediol
- AU Chbihi, M. El M.; Takky, D.; Hahn, F.; Huser, H.; Leger, J. M.; Lamy, C.
- CS Departement de Chimie, Faculte des Sciences Ben M'sik, Universite Hassan II, Casablanca, Morocco
- SO Journal of Electroanalytical Chemistry (1999), 463(1), 63-71 CODEN: JECHES; ISSN: 0368-1874
- PB Elsevier Science S.A.
- DT Journal
- LA English
- AB The electrooxidn. of 1,3-propanediol was studied in acid and alkaline media on platinum and gold electrodes. It is shown by voltammetric techniques that the oxidation of 1,3-propanediol is partially a diffusion controlled. Electrolysis results and in situ FTIR reflectance spectroscopy allowed the authors to identify the adsorbed species and the reaction products. A possible reaction mechanism is discussed.
- RE.CNT 25 THERE ARE 25 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L5 ANSWER 4 OF 4 CAPLUS COPYRIGHT 2007 ACS on STN
- AN 1998:459783 CAPLUS
- DN 129:122962
- TI Cobalt-catalyzed process for preparing 1,3-propanediol
- IN Powell, Joseph Broun; Slaugh, Lynn Henry; Forschner, Thomas Clayton; Lin, Jiang-jen; Thomason, Terry Blane; Weider, Paul Richard; Semple, Thomas Carl; Arhancet, Juan Pedro; Fong, Howard Lam-ho; Mullin, Stephen Blake; Allen, Kevin Dale; Eubanks, David Cleve; Johnson, David William
- PA Shell Oil Co., USA
- SO U.S., 8 pp., Cont.-in-part of U. S. Ser. No. 316,676. CODEN: USXXAM
- DT Patent
- LA English
- FAN.CNT 9

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US S	5981808	Α	19991109	US	1997-	8928	31		199	70715	
PRAI US 1	1994-316660	A2	19940930								
US 1	1994-316669	A2	19940930								
US 1	1994-316676	A2	19940930								
US 1	1994-316680	A2	19940930								
US 1	1996-720270	B1	19960926								
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AB 1,3-Propanediol is prepared in a process which involves hydroformylating ethylene oxide: (a) in an essentially non-water-miscible solvent in the presence of a non-ligated cobalt catalyst and a catalyst promoter at a temperature within the range of about 50° to about 100° C and a pressure within the range of about 500 to about 5000 psig, to produce an intermediate product mixture comprising less than about 15 wt % 3-hydroxypropanal; (b) adding an aqueous liquid and extracting at a temperature less than

about 100° C the 3-hydroxypropanal to provide an aqueous phase comprising 3-hydroxypropanal in greater concentration than the concentration of 3-hydroxypropanal in said intermediate product mixture, and an organic phase comprising the cobalt catalyst; (c) separating the aqueous phase from the organic

phase; (d) hydrogenating the 3-hydroxypropanal to provide a hydrogenation product mixture comprising 1,3-propanediol; and (e) recovering 1,3-propanediol from said hydrogenation product mixture. The process enables the production of 1,3-propanediol in high yields and selectivity without the use of a phosphine ligand-modified cobalt catalyst. Thus, the hydroformylation of ethylene oxide in the presence of dicobaltoctacarbonyl catalyst and NaOAc promoter proceeded at a rate 3-times higher than hydroformylation in the absence of NaOAc.

RE.CNT 28 THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

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L9
     ANSWER 1 OF 7 CAPLUS COPYRIGHT 2007 ACS on STN
     2006:690086 CAPLUS
AN
DN
     145:210617
TI
     Process for preparation of 3-hydroxypropanal and 1,3-propanediol
IN
     Lu, Shunfeng; Wang, Shiliang; Peng, Bin; Qin, Yanhuang; Zhou, Zhiquan; Wu,
PA
     Sinopec Corp., Peop. Rep. China
     Faming Zhuanli Shenqing Gongkai Shuomingshu, 10 pp.
SO
     CODEN: CNXXEV
DT
     Patent
LA
    Chinese
FAN.CNT 1
                 KIND DATE APPLICATION NO. DATE
    PATENT NO.
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    CN 1690033
                              20051102 CN 2004-10037658
PΙ
                        Α
                                                               20040429
PRAI CN 2004-10037658
                              20040429
    This invention provides a process for preparing 3-hydroxypropanal and
     1,3-propanediol comprising reacting epoxyethane with synthetic gas
     followed by hydrogenation. For example, epoxyethane was reacted
     with synthetic gas in the presence of Co2(CO)8 to give 3-hydroxypropanal.
     The 3-hydroxypropanal was hydrogenated in water in the presence
     of Raney Ni to give 1,3-propanediol in high yield.
L9
    ANSWER 2 OF 7 CAPLUS COPYRIGHT 2007 ACS on STN
AN
    2002:695924 CAPLUS
DN
    Process for producing 1,3-propanediol through acrolein hydration and
TΤ
    hydroxypropanal hydrogenation and active hydration catalysts for
     use in the process
IN
    Tsunoda, Takashi; Nomura, Kouji
PA
    Asahi Kasei Kabushiki Kaisha, Japan
SO
    PCT Int. Appl., 35 pp.
    CODEN: PIXXD2
DT
    Patent
LΑ
    Japanese
FAN.CNT 1
                      KIND DATE APPLICATION NO.
    PATENT NO.
                                                              DATE
    WO 2002070447
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                             20020912 WO 2002-JP1921
PΙ
                       A1
                                                                 20020301
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
            CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
            GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KR, KZ, LC, LK, LR, LS,
            LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL,
            PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA,
            UG, US, UZ, VN, YU, ZA, ZM, ZW
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH,
            CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
            BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
    CN 1494523
                        Α
                              20040505
                                        CN 2002-805858 20020301
    TW 224591
                                          TW 2002-91103840
                        В
                               20041201
                                                                 20020301
    US 2004097764
                                          US 2003-469513
                        A1
                               20040520
                                                                 20030902
    US 6911566
                        B2
                               20050628
PRAI JP 2001-57877
                        Α
                               20010302
    JP 2001-75408
                        Α
                               20010316
    JP 2001-389260
                        Α
                               20011221
    WO 2002-JP1921
                        W
                               20020301
AB
    The process comprises hydrating acrolein in a liquid phase in the presence
    of a hydration catalyst to form 3-hydroxypropanal, subsequently removing
    the acrolein when it remains unreacted, and then catalytically
    hydrogenating the 3-hydroxypropanal in a liquid or vapor phase using
    a hydrogenation catalyst, where the hydration catalyst is a catalyst which comprises (a) metalloaluminophosphate type mol. sieves, (b)
    FER zeolites or/and (c) oxides or composite oxides of Groups 4, 13 or/and
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14 elements (provided that crystalline aluminosilicate zeolite is excluded) and which, when dispersed into a 5-fold amount by weight of water, gives a slurry having a pH at 20° of 6 or lower. Thus, mixing water 146.7 with a 85% H3PO4 aqueous solution 95.9, Al(OPr-i)3 169.6, silica powder 2.6 and 20% tetraethylammonium hydroxide aqueous solution 305.2 g at 150° for 133 h, filtering, washing, drying and baking at 500° for 2 h gave a powder having (P2O5 + Al2O3)/SiO2 molar ratio 10:1. Mixing the crystalline powder with 1N NH4NO3 aqueous solution (10% slurry) at room temperature for 3 h,

filtering, washing, drying and baking at 530° for 3 h gave a H-type SAPO-34 catalyst having solution pH (10 g in 50 g water) at 20° of 4.3. Hydration of acrolein using the catalyst had good conversion (49%) and selectivity (96 mol%).

RE.CNT 13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L9 ANSWER 3 OF 7 CAPLUS COPYRIGHT 2007 ACS on STN
- AN 2001:157211 CAPLUS
- DN 135:19301
- TI Modeling the reaction behavior of glycerol in sub- and super-critical water
- AU Anon.
- CS Germany
- SO Wissenschaftliche Berichte Forschungszentrum Karlsruhe (2000), FZKA 6553, 1-187 CODEN: WBFKF5; ISSN: 0947-8620
- DT Report
- LA German
- AB The intention of this thesis was to develop a reaction model to describe the reaction dynamics of glycerol in sub- and supercrit. water (T > 374°, p > 221 bar). The model aimed at improving the understanding of the competition of ionic and radical reactions in this solvent. The model is based on the results of 66 expts. performed in a tube reactor. The reactions were carried out at variable temperature (622-748 K), pressure (250, 350 and 450 bar), reaction time u(32-60 s.), and initial glycerol concentration (1-3 volume%). The product samples consisted of a liquid and a gaseous

phase. A special methodol. for the anal. of each phase had to be worked out. The liquid products were analyzed by headspace-GC-FID and UV-VIS spectroscopy, the gaseous products by GC-WLD-FID. In subcrit. water (622.4 K, 450 bar), mainly the formation of acetaldehyde, acrolein, and formaldehyde could be observed, whereas the formation of gaseous products was negligible. Due to the occurrence of a small number of products only, a high dielec. constant ($\varepsilon = 17$) as well as the high values for the ionic product of water ($Kw = 10-11.2 \text{ mol}_2/I_2$), it was strongly justified to assume a certain ionic reaction mechanism. In supercrit. water, on the other hand, the formation of oxidation and reduction products as a result of reactions with OH and H radicals was observed The following main products were detected as part of the liquid phase: acetaldehyde, acrolein, H2CO, MeOH, allyl alc., propionaldehyde, EtOH, and finally Me2CO. Within the gaseous phase large amts. of CO, CO2, and H2 could be measured. The model calcns. of the reactant and product dynamics were executed by the Chemkin II code on the basis of defined ionic and radical reaction steps. A plug flow behavior under isothermal and isobaric conditions was assumed for the calcns. Due to the lack of well-known reactions in high-temperature water,

both

the chemical and the kinetic parameters, especially for the ionic reactions, were

often highly speculative. Radical reaction steps were based on the classical steps of pyrolysis, whereas the ionic reactions mainly considered bimol. and monomol. elimination as well as aldol condensation and acetalization. The concentration of H+ and OH- was controlled by the self-dissociation of water. The dynamics of reactions and products was optimized stepwise for a pressure of 450 bar. By the results of flow and

sensitivity calcns., a specific variation of the approximated kinetic parameters became possible. These calcns. gave an addnl. hint as to the missing key reactions. In consideration of concerted reaction steps the number of key reactions was substantially reduced and the optimization process simplified. O-containing substances were stabilized by a H-bonding correction. At the end of the optimization process the reaction model consisted of 95 substances, 340 radical and 43 ionic reactions. But only 36 radical and 12 ionic reactions were detected as sensitive key reactions. Exptl. and calculated data differed at lower pressures (250 and 350 bar) as a result of the cage effect. Accordingly, reactions were accelerated with decreasing pressure and a transition from the diffusion controlled reaction to the kinetically controlled reaction was possible. The order n of the global reaction velocity was determined by n = 0.95-1.25. A definite dependence neither on pressure nor on temperature was found. The diagram of Arrhenius showed an overlay of the 2 mechanisms. Whereas the free radical mechanism depended mainly on the temperature, the ionic mechanism was determined by the ionic product of water. An activation energy of 150.3 kJ/mol and a pre-exponential factor of 1018 s-1 could be evaluated. Furthermore, a decline in temperature from 664 K (450 bar) to 622 K (450 bar) was followed by an obvious increase of the global reaction velocity. The pressure rise accelerated the reaction velocity at a temperature of 665-668 K. Between 250-350 bar an activation volume of -167 cm3/mol was estimated Due to the mechanism overlay, the diagram of Kirkwood ($\varepsilon = 2.7-12.3$) also did not show any linear dependency. The global reaction velocity increased with the dielec. constant of the solvent because the transition states had a higher polarity than the reactants. The strongest increase could be observed in the area of the ionic mechanism. Although the model calcns. were conducted on the basis of many assumptions, the results led to a good description of the chemical expts. and, therefore, contributed essentially to the understanding of such a complex reaction system in high-pressure water.

RE.CNT 91 THERE ARE 91 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

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L9 ANSWER 4 OF 7 CAPLUS COPYRIGHT 2007 ACS on STN
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AN 1997:719688 CAPLUS

DN 127:347899

TI Preparation of 1,3-propanediol from ethylene oxide

IN Weider, Paul Richard; Powell, Joseph Broun; Lam, Khiet Thanh

PA Shell Oil Co., USA

SO U.S., 8 pp., Cont.-in-part of U.S. Ser. No. 316,676, abandoned. CODEN: USXXAM

DT Patent

LA English

FAN.CNT 9

	PAT	CENT :	NO.			KIN)	DATE		1	APPL	ICAT	ION I	NO.		D	ATE	
DT						Α	_	1007	1104	1		005					0051	
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	CA	2235	065			С		2006	0103									
	WO	9716	250			A1		1997	0509	1	WO 1	996-	EP47	00		1:	9961	028
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			LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	ΝZ,	PL,	PT,
			RO,	RU,	SD,	SE,	SG,	SI,	SK,	TJ,	TM,	TR,	TT,	UA,	UG,	UΖ,	VN	
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			ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI				
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	ΑU	7000	53			B2		1998	1217									
	EΡ	8624	95			A1		1998	0909]	EP 1	996-	9372	85		1:	9961	028
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	CN	1090	059			В		2002	0904									

	BR	9611339	Α	19990406	BR	1996-	11339		19961	028
	ES	2134016	Т3	19990916	ES	1996-	937285		19961	028
	JP	11515021	T	19991221	JP	1997-	517063		19961	028
	JP	3844781	B2	20061115						
	US	5981808	A	19991109	US	1997-	892831		19970	715
	NO	9801937	Α	19980429	NO	1998-	1937		19980	429
	NO	308894	B1	20001113						
PRAI	US	1994-316676	B2	19940930						
	US	1995-550589	Α	19951031						
	US	1996-720270	B1	19960926						
	WO	1996-EP4700	W	19961028						
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AB 1,3-Propanediol is prepared by: (a) reacting a cobalt salt selected from at least one of cobalt hydroxide, cobalt(II, III) oxide and cobalt carbonate with synthesis gas in a nonwater-miscible liquid to produce a cobalt carbonyl hydroformylation catalyst; (b) hydroformylating ethylene oxide with synthesis gas in a nonwater-miscible liquid medium in the presence of a catalytic amount of the cobalt carbonyl catalyst mixture and a catalyst promoter to produce a product mixture comprising <15% 3-hydroxypropanal; (c) adding an aqueous liquid to the intermediate product mixture and extracting into the aqueous liquid a major portion of the 3-hydroxypropanal

so as to provide an aqueous phase comprising 3-hydroxypropanal in greater concentration than the concentration of 3-hydroxypropanal in the intermediate product

mixture, and an organic phase comprising a major portion of the cobalt carbonyl; (d) separating the aqueous phase from the organic phase; (e) returning a

major portion of the organic phase to step (b); (f) hydrogenating the aqueous phase comprising 3-hydroxypropanal in the presence of a hydrogenation catalyst; and (g) recovering the 1,3-propanediol. A process flow diagram is presented.

- L9 ANSWER 5 OF 7 CAPLUS COPYRIGHT 2007 ACS on STN
- AN 1997:414046 CAPLUS
- DN 127:36203
- TI Preparing 1,3-propanediol
- IN Lam, Khieth Thanh; Powell, Joseph Broun; Weider, Paul Richard
- PA Shell Internationale Research Maatschappij BV, Neth.; Shell Canada Limited
- SO PCT Int. Appl., 23 pp.
 - CODEN: PIXXD2
- DT Patent
- LA English
- FAN.CNT 9

	PATENT NO.								APPLICATION NO.						DATE		
ΡI										 WO 1	 996-1	 EP47	00		19	 9961	028
		AL,															
		DK,	ΕĒ,	ES,	FI,	GB,	GE,	HU,	IL,	IS,	JP,	ΚE,	KG,	ΚP,	KR,	ΚZ,	LC,
		LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	ΝZ,	PL,	PT,
		RO,	RU,	SD,	SE,	SG,	SI,	SK,	TJ,	TM,	TR,	TT,	UA,	ŪĠ,	UΖ,	VN	
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	CA 2235																
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	EP 8624							0909		EP 1	996-	9372	85		19	9961	028
	EP 8624						1999				_						
		BE,	CH,	DE,	•			•	•	•							
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	JP 1151							1221		JP 1	997-	5170	53		19	9961	028
	JP 3844							1115									
	NO 9801	937			Α		1998	0429]	NO 1	998-	1937			19	99804	429

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NO 308894 B1 20001113
PRAI US 1995-550589 A 19951031
US 1994-316676 B2 19940930
WO 1996-EP4700 W 19961028
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AB 1,3-Propanediol is manufactured by (a) reacting ≥1 Co hydroxide
, Co (II, III) oxide, and Co carbonate with synthesis gas in an
essentially non-water-miscible liquid medium under conditions effective to
produce a cobalt carbonyl reaction product comprising at least one active
cobalt carbonyl hydroformylation catalyst species; (b) contacting ethylene
oxide with synthesis gas in an essentially non-water-miscible liquid medium
in the presence of catalytic amount of the cobalt carbonyl reaction product
mixture and an effective amount of a catalyst promoter under reaction
conditions effective to produce an intermediate product mixture comprising
<15% 3-hydroxypropanal (I); (c) adding an aqueous liquid to said intermediate
product mixture and extracting into said aqueous liquid a major portion of the

to provide an aqueous phase containing I in greater concentration than the concentration of I in

the intermediate product mixture and an organic phase containing a major portion of

the cobalt carbonyl; (d) separating the aqueous phase from the organic phase; (e)

returning at least a major portion of the organic phase to the process of step (b); and (f) contacting the aqueous phase containing I with H in the presence

of a hydrogenation catalyst.

- L9 ANSWER 6 OF 7 CAPLUS COPYRIGHT 2007 ACS on STN
- AN 1996:446512 CAPLUS
- DN 125:114171
- TI Process and catalysts for preparing 1,3-alkanediols and 3-hydroxyaldehydes by the hydroformylation of epoxides
- IN Arhancet, Juan Pedro; Forschner, Thomas Clayton; Powell, Joseph Brown; Semple, Thomas Carl; Slaugh, Lynn Henry; Thomason, Terry Blane; Weider, Paul Richard
- PA Shell Internationale Research Maatschappij BV, Neth.; Shell Canada Limited
- SO PCT Int. Appl., 25 pp. CODEN: PIXXD2
- DT Patent
- LA English
- FAN.CNT 2

	PAT	FENT	NO.			KIN	D	DATE			APPL	ICAT	ION :	NO.		D	ATE	
ΡI	WO	9610	550	\supset		A1	-	1996	0411	,	WO 1	 995-:	 EP38	68		1:	9950	928
		₩:-	-AM,	ΑT,	AU,	BB,	ВG,	BR,	BY,	CA,	CH,	CN,	CZ,	DΕ,	DK,	EE,	ES,	FI,
			GB,	GE,	HU,	ıs,	JP,	KΕ,	KG,	ΚP,	KR,	KZ,	LK,	LR,	LT,	LU,	LV,	MD,
			MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,
			TJ,	TM														
		RW:	KE,	MW,	SD,	SZ,	UG,	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙE,	IT,
			LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	ML,	MR,	NE,
			SN,	TD.	TG													
	US	5463	145	ノ)	Α		1995	1031	1	US 1	994-	3166	73		1:	9940	930
	US	-5545	765			Α		1996	0813	•	US 1	994-	3166	61		1:	9940	930
	IN	1921	84			A1		2004	0313		IN 1	995-1	DE17	30		1:	9950	920
	CA	2201	305			A1		1996	0411		CA 1	995-	2201	305		1:	9950	928
	AU	9536	993			A		1996	0426		AU 1	995-	3699	3		1:	9950	928
	AU	6899	63			B2		1998	0409									
	ZA	9508	160			A		1996	0508		ZA 1	995-	8160			1:	950	928
	EP	7834	75			A1		1997	0716		EP 1	995-	9346	52		1:	9950	928
	EP	7834	75			В1		1998	1209									
		R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	IT,	LI,	NL,	PT,	SE			
	CN	1161				Α		1997								1:	9950	928
	CN	1089	083			В		2002	0814									
	BR	9509	054			Α		1998	0623]	BR 1:	995-	9054			19	9950	928

	JP 10507165	T	19980714	JP 1996-511401	19950928
	JP 3779729	B2	20060531		
	RU 2149156	C1	20000520	RU 1997-106785	19950928
	SK 281399	B6	20010312	SK 1997-389	19950928
	PL 181433	B1	20010731	PL 1995-319438	19950928
	NO 9701420	Α	19970325	NO 1997-1420	19970325
	NO 307561	B1	20000425		
	FI 9701267	Α	19970326	FI 1997-1267	19970326
PRAI	US 1994-316661	Α	19940930		
	US 1994-316673	Α	19940930		
	WO 1995-EP3868	W	19950928		
OS	MARPAT 125:114171				

AB The title process comprises hydroformylating an oxirane (e.g., ethylene oxide) with CO and H2 in the presence of Co-based catalyst and a promoter that is a lipophilic quaternary salt of a Group V element [e.g., Bu3(PhCH2)NOAc] to produce 1,3-alkanediols (e.g., 1,3-propanediol) and 3-hydroxyaldehydes (e.g., 3-hydroxypropanal). A process flow diagram is presented.

- L9 ANSWER 7 OF 7 CAPLUS COPYRIGHT 2007 ACS on STN
- 1995:965003 CAPLUS AN
- DN124:88122
- ΤI Cobalt-catalyzed process for manufacturing 1,3-propanediol in high yields without the use of a phosphine-ligated cobalt catalyst
- ΙN Powell, Joseph B.; Slaugh, Lynn H.; Forschner, Thomas C.; Thomason, Terry B.; Semple, Thomas C.; Weider, Paul R.; Arhancet, Juan P.
- PA Shell Oil Co., USA
- U.S., 7 pp. SO
- CODEN: USXXAM
- DTPatent
- T.A English
- FAN.CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI (US 5463144)	A	19951031	US 1994-316671	19940930
PRAI US 1994-316671		19940930		

OS MARPAT 124:88122

AB 1,3-Propanediol is prepared in a process comprising the steps (a) contacting, in an essentially non-water-miscible organic solvent, ethylene oxide with CO and H in the presence of non-phosphine-ligated cobalt compound and an effective amount of a lipophilic quaternary ammonium salt promoter at 50-100° and 500-5000 psig, (b) adding aqueous liquid to intermediate product mixture of (a) comprising <15% 3-hydroxypropanal (I) and extracting into

said aqueous liquid a major portion of the I at <100°, (c) separating the aqueous

phase containing high concentration of I from the organic phase containing catalyst, (d)

contacting the aqueous phase with H in the presence of a hydrogenation catalyst at ≥100 psig and ≥40° to provide a hydrogenation product mixture comprising 1,3-propanediol, (e) recovering 1,3-propanediol from the hydrogenation product mixture; and (f) returning at least a portion of the organic phase comprising cobalt compound and lipophilic quaternary ammonium salt to the process of step (a). Thus, 0.46 g of 40% benzyltrimethylammonium methoxide promoter in MeOH, was added in the hydroformylation reaction and 4.73% I was formed at a rate of 11.8 h-1, or a 2.6-fold rate increase over that observed in the absence of promoter.